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**UNITED STATES DISTRICT COURT
CENTRAL DISTRICT OF CALIFORNIA**

TAYLOR DEVORAK,

Plaintiff,

vs.

**PFIZER INC.; VIATRIS INC.;
GREENSTONE LLC; PRASCO, LLC d/b/a
PRASCO LABS.; PHARMACIA &
UPJOHN CO. LLC; and PHARMACIA
LLC,**

Defendants.

**COMPLAINT AND DEMAND
FOR JURY TRIAL**

Case No.: 5:24-cv-02349

Plaintiff Taylor Devorak, by and through Plaintiff's undersigned counsel, brings this civil action against Defendants for personal injuries and damages suffered by Plaintiff, and alleges upon information and belief as follows:

INTRODUCTION

1. This is an action for damages related to Defendants' wrongful conduct in connection with the development, design, testing, manufacturing, labeling, packaging, promoting, advertising, marketing, distribution, and selling of medroxyprogesterone acetate (hereinafter "MPA"), also known as depot medroxyprogesterone acetate (hereinafter "DMPA"). Defendants' trade name for this prescription drug is Depo-Provera[®] (hereinafter "Depo-Provera").

1 2. Defendants manufacture, promote, and sell Depo-Provera as a prescription drug used
2 for contraception or to treat endometriosis, among other indications. Depo-Provera is manufactured as
3 an injection to be administered intramuscularly every three (3) months in either the upper arm or
4 buttocks.

5 3. Depo-Provera injured Plaintiff Taylor Devorak (hereinafter “Plaintiff”) by causing
6 or substantially contributing to the development of an intracranial meningioma, a type of brain tumor,
7 which has caused serious injuries.

8 4. Defendants knew or should have known for decades that Depo-Provera, when
9 administered and prescribed as intended, can cause or substantially contribute to the development of
10 meningiomas.

11 5. Several scientific studies have established that progesterone, its synthetic analogue
12 progestin, and Depo-Provera in particular, cause or substantially contribute to the development of
13 intracranial meningioma, a type of brain tumor.

14 6. Nevertheless, Defendants failed to warn, instruct, advise, educate, or otherwise
15 inform Depo-Provera users and prescribers about the risk of intracranial meningioma or the need for
16 monitoring for resultant symptoms.

17 7. To date, the U.S. label for Depo-Provera still makes no mention of the increased risk
18 to patients of developing intracranial meningiomas despite the fact that the European Union (EU) and
19 the United Kingdom labels now list meningioma under the “special warnings and precautions for use”
20 section and advise EU patients to speak with their doctors before using Depo-Provera if they have any
21 history of meningioma.

22 8. Moreover, the Canadian label for Depo-Provera has listed “meningioma” among its
23 “Post-Market Adverse Drug Reactions” since at least 2015.
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1 19. Prasco has a registered agent for service of process, CT Corp., at 330 North Brand
2 Boulevard in Glendale, CA.

3 20. Defendant PHARMACIA & UPJOHN CO. LLC (hereinafter “Pharmacia & Upjohn”
4 or “Upjohn”) is or was a corporation organized under Michigan law and headquartered at 7171 Portage
5 Road, Kalamazoo, MI 49002.

6 21. Pharmacia & Upjohn has a registered agent for service of process, CT Corp., at 330
7 North Brand Boulevard in Glendale, CA.

8 22. Defendant PHARMACIA LLC (hereinafter “Pharmacia”) is a corporation organized
9 under Delaware law and headquartered at Pfizer Peapack Campus, 100 Route 206 North, Peapack, NJ
10 07977.

11 23. Pharmacia has a registered agent for service of process, CT Corp., at 820 Bear Tavern
12 Road, West Trenton, NJ 08628.

13 24. Defendant Pfizer is the current New Drug Application (hereinafter “NDA”) holder
14 for Depo-Provera and has solely held the NDA for Depo-Provera since 2020. Upon information and
15 belief, Pfizer has effectively held the NDA since at least 2002 when it acquired Pharmacia & Upjohn—
16 who then held the NDA—as a wholly owned subsidiary. No later than 2003 did Pfizer’s name appear
17 on the label alongside Pharmacia & Upjohn.

18 25. At all relevant times, Defendant Pharmacia & Upjohn was a wholly owned
19 subsidiary of Defendant Pfizer until Upjohn was spun off in a merger in 2020 to create Defendant
20 Viartis and the remnant, i.e., Defendant Pharmacia, was retained by Pfizer.

21 26. Defendant Greenstone, founded in 1993, was a wholly owned subsidiary of Pfizer,
22 that at pertinent times was in the business of offering a product portfolio of “authorized generic”
23 medicines, including Depo-Provera.
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1 27. Defendant Greenstone is a company that until November 2020 was styled as a wholly
2 owned subsidiary of Pfizer but was in fact exclusively staffed with Pfizer personnel who reported to
3 Pfizer's HR department, were on Pfizer's payroll, and shared the same corporate space with Pfizer in
4 Peapack, NJ. Pfizer also managed Greenstone's key business functions including financial and sales
5 analysis, business technology, customer service, legal matters, intellectual property, and supply chain
6 operations. Thus, Greenstone was effectively a department within Pfizer.

7 28. Defendants Greenstone/Pfizer sold a "generic" version of Depo-Provera that was in
8 fact what is known as an "authorized generic." Unlike standard generics, which must contain only the
9 same active ingredients and have the same pharmaceutical effect but can otherwise contain vastly
10 different additives, "authorized generics" are exact replicas of the brand name drug, with the identical
11 chemical composition, simply marketed without the brand-name on its label. In other words,
12 Greenstone was presenting itself as a distinct generic manufacturing entity when it was in fact Pfizer
13 personnel producing the exact same brand-name Depo-Provera at Pfizer's own facility.

14 29. The FDA has stated that the term "authorized generic" drug is most commonly used
15 to describe an approved brand name drug that is marketed without the brand name on its label. Other
16 than the fact that it does not have the brand name on its label, it is the exact same drug product as the
17 branded product. An "authorized generic" may be marketed by the brand name drug company, or
18 another company with the brand company's permission.¹

19 30. Indeed, Pfizer's own website still states that "GREENSTONE Authorized Generics
20 are manufactured to the same standards and at the same facilities as Pfizer brand-name drugs."²
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26 ¹ See [https://www.fda.gov/drugs/abbreviated-new-drug-application-anda/fda-list-authorized-generic-](https://www.fda.gov/drugs/abbreviated-new-drug-application-anda/fda-list-authorized-generic-drugs)
27 [drugs](https://www.fda.gov/drugs/abbreviated-new-drug-application-anda/fda-list-authorized-generic-drugs) (last accessed Sept. 30, 2024).

28 ² See [https://www.pfizer.com/news/press-release/press-release-detail/pfizers-greenstone-and-digital-](https://www.pfizer.com/news/press-release/press-release-detail/pfizers-greenstone-and-digital-mens-health-clinic-roman)
[mens-health-clinic-roman](https://www.pfizer.com/news/press-release/press-release-detail/pfizers-greenstone-and-digital-mens-health-clinic-roman) (last accessed Sept. 26, 2024).

1 31. Pfizer was the actual manufacturer of the authorized generic product that Greenstone
2 distributed and sold.

3 32. Defendant Viartis was formed by the merger of Upjohn, Greenstone, and another
4 company, Mylan N.V., in November 2020. Viartis is thus merely the latest iteration of Upjohn and
5 Greenstone.

6 33. Even after the merger, Defendant Greenstone has continued to operate from the same
7 location at Pfizer's corporate offices in Peapack, NJ.

8 34. Additionally, Defendant Pfizer retained 57% ownership of Viartis stock, making
9 Pfizer the majority owner of Viartis, and since Pfizer retained the remnants of Pharmacia, Pfizer
10 effectively remains the majority owner of Defendants Pharmacia & Upjohn and Greenstone.
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12 35. Defendant Prasco is another "authorized generic" manufacturer of Depo-Provera,
13 meaning Prasco simply takes brand-name Depo-Provera manufactured by Defendants Greenstone
14 and/or Pfizer and distributes it as its own generic product.
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16 36. Defendant Prasco consistently maintains a sizeable percentage of the market share
17 for Depo-Provera sales in the US.

18 37. Pfizer is the actual manufacturer of the authorized generic product that Prasco
19 distributes and sells. Pfizer packages and labels the product with the Prasco name on the label under
20 the Pfizer NDA.

21 38. All Defendants do business in California by, among other things, distributing,
22 marketing, selling, and/or profiting from brand name and/or "authorized generic" Depo-Provera in
23 California, as well as throughout the United States.
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25 39. At all times material herein, Defendants were, and still are, pharmaceutical companies
26 involved in the manufacturing, research, development, marketing, distribution, sale, and release for
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1 use to the general public of pharmaceuticals, including Depo-Provera and its “authorized generic”
2 version, in California, and throughout the United States.

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4 **JURISDICTION AND VENUE**

5 40. This Court has diversity jurisdiction over this action pursuant to 28 U.S.C. § 1332,
6 as the amount in controversy exceeds \$75,000.00 and the Parties are citizens of different States.

7 41. All Defendants regularly conduct business in California.

8 42. This Court has supplemental jurisdiction over the remaining common law and state
9 claims pursuant to 28 U.S.C. § 1367.

10 43. Venue is proper in this Court pursuant to 28 U.S.C. § 1391 because a substantial part
11 of the events or omissions giving rise to the claim, including the distribution, sale, and administration
12 of Depo-Provera to Plaintiff and Plaintiff’s development, diagnosis, and treatment of meningioma, all
13 occurred in the Central District of California.

14 44. Defendant Pfizer has extensive connections to the State of California that are highly
15 relevant to the subject matter of the instant action.

16 45. For example, Pfizer maintains the Pfizer La Jolla Research Site, a 25-acre “campus”
17 complete with a 500,000-square-foot state-of-the-art facility devoted to the study of oncology, drug
18 safety, and pharmacokinetics.³

19 46. As of December 2018, Defendant Pfizer’s La Jolla campus is home to more than 900
20 scientists and clinicians studying, *inter alia*, the effects of drugs on the development of tumors.⁴

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27 ³ <https://www.pfizer.com/la-jolla-california> (Last accessed Oct. 13, 2024).

28 ⁴ See <https://www.sandiegouniontribune.com/2018/12/11/pfizer-adds-100-to-cancer-research-center-in-la-jolla/> (Dec. 11, 2018) (Last accessed Oct. 13, 2024).

1 47. According to Pfizer’s website, the “Pfizer La Jolla campus is an important part of
2 California’s life sciences community and partners with academic institutions and other research
3 organizations to advance scientific understanding and deliver new medicines.”⁵

4 48. Pfizer’s website states: “In 2011, Pfizer announced that it is partnering with the
5 University of California, San Diego Health Sciences and Sanford-Burnham Medical Research Institute
6 through [Pfizer’s] Centers for Therapeutic Innovation (CTI).” Pfizer’s website explains “CTI is a
7 network of collaborative partnerships with top-tier life science research institutions in California,
8 Massachusetts and New York that aims to accelerate and transform drug discovery and development.
9 In San Diego, CTI’s home base is located on the Pfizer La Jolla campus.”⁶

10 49. CTI was launched by Pfizer in 2010 as “an entrepreneurial network of partnerships
11 with leading academic medical centers to transform research and development by accessing leading
12 translational researchers.”⁷

13 50. The University of California, San Francisco was “the first collaboration in the
14 network.”⁸

15 51. Pfizer’s senior vice president of Worldwide BioTherapeutics Research and
16 Development stated at the time of the announcement, “UCSF is a world-class academic medical center
17 with a strong focus on both basic science and clinical research, which is why Pfizer is partnering with
18 them on this initiative. Ultimately, we believe this could create significant benefit for the patient.”⁹

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24 ⁵ <https://www.pfizer.com/la-jolla-california> (Last accessed Oct. 13, 2024).

25 ⁶ *Id.*

26 ⁷ https://www.pfizer.com/news/press-release/press-release-detail/pfizer_launches_global_centers_for_therapeutic_innovation_a_network_of_research_partners_hips_with_university_of_california_san_francisco (Nov. 16, 2010) (Last accessed Oct. 13, 2024).

27 ⁸ *Id.*

28 ⁹ *Id.*

1 52. Pfizer has thus deliberately created strong connections not just to the consumers and
2 patients of California but also to the life and health sciences communities and the State educational
3 institutions of California as well.

4 53. Moreover, Defendants Pfizer, Viatris, Upjohn & Pharmacia, and Prasco are all
5 registered to do business in the State of California and can be served at their registered agent for service
6 of process, CT Corp., at 330 North Brand Boulevard in Glendale, CA.

7 54. All Defendants at different periods of time had a contractual and/or sales relationship
8 directly or through intermediaries to sell Depo-Provera to Kaiser Permanente Health System knowing
9 that health care providers at Kaiser Permanente in California would be injecting Depo-Provera into
10 patients.

11 55. At various points of time, Defendant Pfizer sponsored continuing education courses,
12 seminars, and meetings to promote the use of Depo-Provera to Plaintiff's health care providers.
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14 **PLAINTIFF TAYLOR DEVORAK'S SPECIFIC FACTS**

15 56. On April 27, 2015, at the age of 20, Plaintiff Taylor Devorak was first administered
16 Depo-Provera as a contraceptive by her healthcare provider at Planned Parenthood in Victorville, CA.
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18 57. At all times relevant herein, Defendants represented Depo-Provera to be appropriate,
19 safe, and suitable for such purposes through the label, packaging, patient inserts, and advertising.

20 58. Between April 27, 2015, and May 5, 2024, Plaintiff received regular Depo-Provera
21 injections as prescribed, totaling approximately thirty-six (36) injections. These injections were
22 administered according to her healthcare provider's recommendations.

23 59. Over time, Plaintiff began to experience troubling symptoms, including blurry vision
24 beginning around August 16, 2018, as well as difficulty focusing on objects with her right eye and
25 noticing double vision. These symptoms significantly impacted her daily activities, including driving
26 and performing her duties as a phlebotomist.
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1 60. On August 16, 2018, Plaintiff underwent a CT scan of her head at Loma Linda
2 University Health, which revealed a 2.5 cm suprasellar mass exerting pressure on the optic chiasm.

3 61. An MRI performed the following day, on August 17, 2018, confirmed a 2.4 x 2.4 x
4 2.2 cm sellar/suprasellar meningioma, specifically located at the planum sphenoidale and compressing
5 the optic chiasm.

6 62. Due to the tumor's impact on her vision and quality of life, Plaintiff underwent
7 intracranial surgery on September 12, 2018, to resect the meningioma through an endoscopic
8 transsphenoidal approach. Her skull base was reconstructed using a vascularized pedicle flap.

9 63. Post-surgery imaging on September 12, 2018, and subsequent follow-up imaging
10 indicated residual tissue along the surgical bed. A CT scan and MRI performed on September 13,
11 2018, identified T2 signal abnormalities and thinning of the optic chiasm.

12 64. Continued MRIs, including those conducted on January 2019, July 1, 2019, and
13 October 29, 2019, indicated an enhancing soft tissue mass along the roof of the sella, measuring
14 approximately 14 x 11 x 4 mm, adhering to the right cisternal optic nerve and the inferior aspect of
15 the optic chiasm, likely indicative of residual meningioma or related tissue.

16 65. Due to the sensitive location of Plaintiff's residual meningioma and the impact on
17 her optic chiasm, her medical team has advised against further invasive procedures, as additional
18 surgery would present significant risks to her vision and health.

19 66. As a result of Defendants' actions and inactions, Plaintiff has suffered serious
20 injuries and damages due to Plaintiff's development of a heavily calcified intracranial meningioma
21 and sequelae related thereto.

22 67. Plaintiff was unaware until very recently, following publicity associated with a large
23 case control study in France published in March 2024, that Depo-Provera had any connection to her
24 meningioma.

GENERAL ALLEGATIONS

A. Intracranial Meningioma

68. Intracranial meningioma is a medical condition in which a tumor forms in the meninges, the membranous layers surrounding the brain and spinal cord.

69. Although the tumor formed by an intracranial meningioma is typically histologically benign (meaning it usually does not metastasize), the growing tumor can nevertheless press against the sensitive surrounding tissues, i.e., the brain, and thereby cause a number of severe and debilitating symptoms ranging from seizures and vision problems to weakness, difficulty speaking, and even death, among others. Moreover, a sizeable number of meningiomas (15-20%) do become metastatic, greatly increasing their danger.

70. Treatment of a symptomatic intracranial meningioma typically requires highly invasive brain surgery that involves the removal of a portion of the skull, i.e., a craniotomy, in order to access the brain and meninges. Radiation therapy and chemotherapy may also be required as the sensitive location of the tumor in the brain can render complete removal highly risky and technically difficult.

71. Due to the sensitive location of an intracranial meningioma immediately proximate to critical neurovascular structures and the cortical area, surgery can have severe neurological consequences. Many studies have described the potential for postoperative anxiety and depression and an attendant high intake of sedatives and antidepressants in the postoperative period. Surgery for intracranial meningioma can also lead to seizures requiring medication to treat epilepsy. Moreover, meningiomas related to progesterone-based contraceptives tend to manifest at the base of the skull where removal is even more challenging, further increasing the risks of injuries.

B. Depo-Provera

1 72. Depo-Provera (depot medroxyprogesterone acetate, hereinafter “DMPA”) was first
2 approved by the FDA in 1992 to be used as a contraceptive, and later, with the approval of the Depo-
3 SubQ Provera 104 variant in 2004, as a treatment for endometriosis.

4 73. Depo-Provera is administered as a contraceptive injection that contains a high dose
5 of progestin, a synthetic progesterone-like hormone that suppresses ovulation.

6 74. According to a recent National Health Statistics Report published in December 2023,
7 nearly a quarter (24.5%) of all sexually experienced women in the United States between 2015 and
8 2019 had ever used Depo-Provera.¹⁰

9 75. According to that same report, those proportions increase even further for Hispanic
10 (27.2%) women and Black (41.2%) women who had ever used Depo-Provera.¹¹

11 76. Depo-Provera is a 150 mg/mL dosage of DMPA that is injected every three (3)
12 months into the deep tissue musculature of either the buttocks or the upper arm, with present labelling
13 recommending alternating the injection site at each injection.

14 77. Defendant Pfizer represents Depo-Provera to be one of the most effective
15 contraceptives in existence. In fact, the Depo-Provera label groups injectable contraceptives like
16 Depo-Provera alongside “Sterilization” as the most effective contraceptive methods resulting in the
17 fewest unintended pregnancies.

18 78. Among reproductive age women who used any form of contraception from 2017-
19 2019, the contraceptive injection was most often used by young women, lower-income women, and
20 Black women.¹²

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26 ¹⁰ Daniels, K et al., “Contraceptive Methods Women Have Ever Used: United States, 2015-2019”,
Nat’l Health Statistics Report, No. 195, Dec. 14, 2023.

27 ¹¹ *Id.*

28 ¹² See <https://www.kff.org/womens-health-policy/fact-sheet/dmpa-contraceptive-injection-use-and-coverage/> (last accessed Sept. 30, 2024).

1 79. Depo-Provera was first developed by Defendant Upjohn (later acquired by
2 Defendant Pfizer) in the 1950s.

3 80. Upjohn introduced Depo-Provera as an injectable intramuscular formulation for the
4 treatment of endometrial and renal cancer in 1960.

5 81. The NDA for Depo-Provera for use as a contraceptive was originally submitted to
6 the FDA by Upjohn in 1967; however, this application was rejected.

7 82. Upjohn again applied to the FDA for approval to market Depo-Provera as a
8 contraceptive in 1978 but was again rebuffed.

9 83. Upjohn applied to the FDA for a third time for the approval of Depo-Provera as a
10 contraceptive in 1983, but the FDA once again rejected the application.

11 84. As early as 1969, Upjohn successfully received approval for Depo-Provera for
12 contraception in international markets, including France.

13 85. Upjohn's NDA for Depo-Provera for use as a contraceptive was eventually approved
14 by the FDA on or about October 29, 1992.

15 86. Upjohn merged with Swedish manufacturer Pharmacia AB to form Pharmacia &
16 Upjohn in 1995.

17 87. Defendant Pfizer acquired Pharmacia & Upjohn in 2002, thereby acquiring the Depo-
18 Provera NDA as well as the associated responsibilities and liabilities stemming from the
19 manufacturing, sale, and marketing of Depo-Provera.

20 88. Pfizer has effectively held the Depo-Provera NDA since acquiring Pharmacia &
21 Upjohn in 2002, and has solely held the NDA since 2020, when Upjohn was spun off to form
22 Defendant Viatrix.

1 89. Throughout the time Defendants marketed both variants of Depo-Provera,
2 Defendants failed to provide adequate warnings to patients and the medical community, including
3 Plaintiff's prescribing physician, of the risks associated with using the drug.

4 90. Defendants also failed to adequately test Depo-Provera to investigate the potential
5 for intracranial meningioma.

6 91. Defendants are also liable for the conduct of its predecessors who failed to adequately
7 design, test, and warn of the dangers associated with use of Depo-Provera.
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9 **C. The Dangers of Depo-Provera**

10 92. The association between progesterone and meningioma has been known or knowable
11 for decades, particularly for sophisticated pharmaceutical corporations like Defendants engaging in
12 FDA-required post-market surveillance of their products for potential safety issues. That duty includes
13 an obligation to keep current with emerging relevant literature and where appropriate, perform their
14 own long- term studies and follow-up research.

15 93. Since at least 1983, the medical and scientific communities have been aware of the
16 high number of progesterone receptors on meningioma cells, especially relative to estrogen
17 receptors.¹³
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19 94. This finding was surprising and notable within the medical and scientific
20 communities because it had previously been thought that meningioma cells, like breast cancer cells,
21 would show a preference for estrogen receptors.¹⁴ Researchers publishing in the *European Journal of*
22 *Cancer and Clinical Oncology* instead found the opposite, indicating progesterone was involved in
23 the incidence, mediation, and growth rate of meningiomas.¹⁵ This particular study was published
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25 ¹³ See Blankenstein, et al., "Presence of progesterone receptors and absence of oestrogen receptors in
26 human intracranial meningioma cytosols," *Eur J Cancer & Clin Oncol*, Vol. 19, No. 3, pp. 365-70
27 (1983).

27 ¹⁴ See *id.*

28 ¹⁵ See *id.*

1 nearly a decade before the FDA approved Depo-Provera for contraception in 1992. In those nine (9)
2 years before Depo-Provera was approved for contraception, and in the thirty-two (32) years since—
3 more than forty (40) years in all—Defendants have seemingly failed to investigate the effect of their
4 high-dose progesterone Depo-Provera on the development of meningioma.

5 95. Since at least as early as 1989, researchers have also been aware of the relationship
6 between progesterone-inhibiting agents and the growth rate of meningioma.¹⁶ That year, the same
7 authors published a study in the *Journal of Steroid Biochemistry* entitled, “Effect of steroids and
8 antisteroids on human meningioma cells in primary culture,” finding that meningioma cell growth was
9 significantly reduced by exposure to mifepristone, an antiprogesterone agent.¹⁷

10 96. Numerous studies published in the decades since have presented similar findings on
11 the negative correlation between progesterone-inhibiting agents and meningioma.¹⁸

12 97. Relatedly, a number of studies published in the interim have reported on the positive
13 correlation between a progesterone and/or progestin medication and the incidence and growth rate of
14 meningioma.¹⁹

15 98. In 2015, a retrospective literature review published in the peer-reviewed journal
16 *BioMed Research International* by Cossu, et al. surveyed the relevant literature including many of the

17 ¹⁶ See Blankenstein, et al., “Effect of steroids and antisteroids on human meningioma cells in primary
18 culture,” *J Steroid Biochem*, Vol. 34, No. 1-6, pp. 419-21 (1989).

19 ¹⁷ See *id.*

20 ¹⁸ See, e.g., Grunberg, et al., “Treatment of unresectable meningiomas with the antiprogesterone agent
21 mifepristone,” *J Neurosurgery*, Vol. 74, No. 6, pp. 861-66 (1991); see also Matsuda, et al., “Antitumor
22 effects of antiprogesterones on human meningioma cells in vitro and in vivo,” *J Neurosurgery*, Vol.
23 80, No. 3, pp. 527-34 (1994).

24 ¹⁹ See, e.g., Gil, et al., “Risk of meningioma among users of high doses of cyproterone acetate as
25 compared with the general population: evidence from a population-based cohort study,” *Br J Clin
26 Pharmacol*. Vol. 72, No. 6, pp. 965-68 (2011); see also Bernat, et al., “Growth stabilization and
27 regression of meningiomas after discontinuation of cyproterone acetate: a case series of 12 patients,”
28 *Acta Neurochir (Wien)*. Vol. 157, No. 10, pp. 1741-46 (2015); see also Kalamarides, et al., “Dramatic
shrinkage with reduced vascularization of large meningiomas after cessation of progestin treatment,”
World Neurosurg. Vol. 101, pp 814.e7-e10 (2017).

1 studies cited above and concluded that mifepristone, an antiprogesterone agent, had a regressive effect
2 on meningioma, meaning it stopped or reversed its growth.²⁰ Reviewing the Blankenstein studies as
3 well as many others conducted over a span of more than thirty (30) years, the authors concluded that
4 mifepristone competes with progesterone for its receptors on meningioma cells and, by blocking
5 progesterone from binding, stems or even reverses the growth of meningioma.

6 99. In light of the aforementioned studies, for several decades the manufacturers and
7 sellers of Depo-Provera and its authorized generic and generic analogues, Defendants, had an
8 unassignable duty to investigate the foreseeable potential that a high dose synthetic progesterone
9 delivered in the deep tissue could cause the development or substantially contribute to the growth of
10 meningioma. Defendants were also best positioned to perform such investigations. Had Defendants
11 done so, they would have discovered decades ago that their high dose progestin Depo-Provera was
12 associated with a highly increased risk of meningioma and would have spared Plaintiff and countless
13 others the pain and suffering associated with meningioma. Instead, Defendants did nothing, and
14 therefore willfully failed to apprise the medical community, and the women patients receiving
15 quarterly high dose injections, of this dangerous risk.

16 100. Indeed, more recently, researchers have found that prolonged use (greater than one
17 year) of progesterone and progestin, and specifically Depo-Provera, is linked to a greater incidence of
18 developing intracranial meningioma, as would be expected based on all the aforementioned studies and
19 recognition of the relationship between dose and duration of use and the development of adverse events
20 well recognized in the fields of pharmacology, toxicology, and medicine.

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27 ²⁰ See Cossu et al., “The Role of Mifepristone in Meningiomas Management: A Systematic Review of
28 the Literature” *BioMed Res. Int.* 267831 (2015), <https://doi.org/10.1155/2015/267831>

1 101. In 2022, an article was published in the journal *Endocrinology* entitled “Estrogen and
2 Progesterone Therapy and Meningiomas.”²¹ This retrospective literature review noted that a “dose-
3 dependent relationship” has been established between at least one progestin and the incidence and
4 growth rate of meningioma. The study authors further noted that progesterone-mediated meningiomas
5 appear to be located most often in the anterior and middle base of the skull and are more likely to be
6 multiple and require more intensive treatment.

7 102. In 2023, researchers reported on a direct link between Depo-Provera and
8 meningioma. That year a case series was published in the *Journal of Neurological Surgery Part B:*
9 *Skull Base* titled “Skull Base Meningiomas as Part of a Novel Meningioma Syndrome Associated with
10 Chronic Depot Medroxyprogesterone Acetate Use.”²² The abstract reported on 25 individuals who
11 developed one or more intracranial meningiomas related to chronic use of Depo-Provera. Of the
12 twenty-five (25) patients, ten (10) were instructed to cease Depo-Provera use, after which five (5) of
13 those patients had “clear evidence of tumor shrinkage,” leading the authors to conclude “there appears
14 to be a clear progestin meningioma syndrome associated with chronic DMPA use.”
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16 103. In 2024, the French National Agency for Medicines and Health Products Safety
17 along with several French neurosurgeons, epidemiologist, clinicians, and researchers published a large
18 case control study in the *British Medical Journal (BMJ)*, one of the premier scientific journals in the
19 world, to assess the risk of intracranial meningioma with the use of numerous progestogens among
20 women in France, hereinafter referred to as the *Roland* study.²³
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24 ²¹ Hage, et al., “Estrogen and progesterone therapy and meningiomas,” *Endocrinology*, Vol. 163, pp.
1-10 (2022).

25 ²² Abou-Al-Shaar, et al., “Skull base meningiomas as part of a novel meningioma syndrome associated
26 with chronic depot medroxyprogesterone acetate use,” *J Neurol Surg Part B Skull Base*, Vol. 84:S1-
344 (2023).

27 ²³ Roland, et al., “Use of progestogens and the risk of intracranial meningioma: national case-control
28 study,” *BMJ*, Vol. 384, published online Mar. 27, 2024 at <https://doi.org/10.1136/bmj-2023-078078>
(last accessed Apr. 21, 2024).

1 104. By way of history, the *Roland* study noted that concerns over meningiomas associated
2 with high dose progestogen medications resulted in the recent discontinuation of three such medications
3 in France and the EU. Specifically, there were “postponements in the prescription of chlormadinone
4 acetate, nomegestrol acetate, and cyproterone acetate, following the French and European
5 recommendations to reduce the risk of meningioma attributable to these progestogens in 2018 and
6 2019.”²⁴

7 105. The study analyzed 18,061 cases of women undergoing surgery for intracranial
8 meningioma between 2009 and 2018. The study found that “prolonged use of ... medroxyprogesterone
9 acetate [Depo-Provera] ... was found to increase the risk of intracranial meningioma.” Specifically,
10 the authors found that prolonged use of Depo-Provera resulted in a 555% increased risk of developing
11 intracranial meningioma. The study authors concluded “[t]he increased risk associated with the use of
12 injectable medroxyprogesterone acetate, a widely used contraceptive,” was an important finding. The
13 authors also noted Depo-Provera is “often administered to vulnerable populations,” i.e., lower-income
14 women who have no other choice but to take the subsidized option which only requires action every
15 three months to remain effective for its intended use of preventing pregnancy, and, in the case of the
16 subcutaneous variant, treating endometriosis.

17 106. The 2024 *Roland* study published in *BMJ* studied the effect of several other
18 progestogen-based medications. Three study subjects showed no excess risk of intracranial
19 meningioma surgery with exposure to oral or intravaginal progesterone or percutaneous progesterone,
20 dydrogesterone or spironolactone, while no conclusions could be drawn for two others due to lack of
21 exposed cases. The other medications, including medroxyprogesterone acetate (Depo-Provera), were
22 found to be associated with an increased risk of intracranial meningioma, with Depo-Provera having
23
24
25
26

27 _____
28 ²⁴ See *id.*

1 by far the second highest increased risk, surpassed only by the product cyproterone acetate, which had
2 already been withdrawn from the market due to its association with meningioma.

3 107. Depo-Provera had by far the highest risk of meningioma surgeries amongst
4 progesterone contraceptive products studied, rendering Depo-Provera more dangerous than other
5 drugs and treatment options designed to prevent pregnancy due to the unreasonably increased risk of
6 injury associated with intracranial meningioma, including but not limited to seizures, vision problems,
7 and even death.

8
9 108. Further, the *Roland* study found the longer duration of exposure had a greater risk
10 noting the results show that three quarters of the women in the case group who had been exposed for
11 more than a year had been exposed for more than three years.

12 109. The *Roland* study noted that among cases of meningioma observed in the study,
13 28.8% (5,202/18,061) of the women used antiepileptic drugs three years after the index date of
14 intracranial surgery.

15
16 110. More recently, in September 2024, an article entitled “The Association between
17 Medroxyprogesterone Acetate Exposure and Meningioma” was published in *Cancers*. This large case-
18 control study analyzed over 117,000 meningioma cases and more than one million matched controls
19 and found that “injection exposure” of medroxyprogesterone acetate, i.e. Depo-Provera usage, was
20 associated with a 53% increase in the development of meningioma. The association was specific to
21 cerebral meningiomas and became even stronger with prolonged use.²⁵

22
23 111. In October 2024, researchers at the University of Cincinnati published an abstract in
24 the *International Journal of Radiation Oncology Biology Physics* titled “Progesterone Contraception
25 and Tumor-Related Visual Impairment in Premenopausal Women with Meningioma Referred for
26

27 ²⁵ Griffin, “The association between medroxyprogesterone acetate exposure and meningioma,”
28 *Cancers*, Vol. 16, No. 3362 (2024).

1 Radiation.” This paper reported on a retrospective case-control study that examined, *inter alia*, the
2 role of hormonal contraception in the development of intracranial meningioma causing visual
3 impairment in women under the age of 55. The authors concluded “progesterone use is a significant
4 risk factor for meningioma-related visual deficits ..., with a disproportionate number on [Depo-]
5 Provera specifically.”²⁶

6 **D. Defendants’ Failure to Test Depo-Provera**

7
8 112. Defendants knew or should have known of the potential impact of the drug to cause the
9 development of intracranial meningioma but failed to adequately study these adverse effects.

10 113. Furthermore, despite the fact that studies have emerged over the course of decades
11 providing evidence of the meningioma-related risks and dangers of progesterone and progestins and
12 Depo-Provera specifically, Defendants have failed to adequately investigate the threat that Depo-Provera
13 poses to patients' well-being or warn the medical community and patients of the risk of intracranial
14 meningioma and sequelae related thereto.

15 **E. Defendants’ Continuing Failure to Disclose Depo-Provera’s Health Risks**

16
17 114. According to the Drugs@FDA website, the label for Depo-Provera has been updated
18 on at least thirteen (13) occasions since 2003, with the most recent update coming in July 2024.²⁷
19 Despite the fact there are at least fourteen (14) iterations of the Depo-Provera label, Defendants’ labels
20 have not contained any warning or any information whatsoever on the increased propensity of Depo-
21 Provera to cause severe and debilitating intracranial meningioma like that suffered by Plaintiff.

22
23
24
25 ²⁶ Bailey, et al., “Progesterone contraception and tumor-related visual impairment in premenopausal
26 women with meningioma referred for radiation,” *Int’l J of Radiation Oncology Biology Physics*, Vol.
120, No. 2 Supp., pp. E217 (2024).

27 ²⁷ See Drugs@FDA:FDA-Approved Drugs- Depo-Provera,
28 <https://www.accessdata.fda.gov/scripts/cder/daf/index.cfm?event=overview.process&ApplNo=020246> (last visited Apr. 29, 2024).

1 115. Despite the aforementioned article in the *BMJ* and all the preceding medical literature
2 cited above demonstrating the biological plausibility of the association between progesterone and
3 meningioma, evidence of Depo-Provera related cases of meningioma and the evidence of other high
4 dose progestones causing meningiomas, Defendants have still made no change to the U.S. Depo-
5 Provera label related to intracranial meningioma. Furthermore, Defendants have failed to take any steps
6 to otherwise warn the medical community and Depo-Provera users of these significant health risks,
7 despite changing the label as recently as July 2024 to include warnings about pregnancy-related risks,
8 and despite Defendant Pfizer stating to The Guardian when the *BMJ* article was released in April 2024:
9 “We are aware of this potential risk associated with long-term use of progestogens and, in collaboration
10 with regulatory agencies, are in the process of updating product labels and patient information leaflets
11 with appropriate wording.”²⁸

13 116. Defendant Pfizer *has* changed the label in the EU and the UK and potentially in other
14 countries. Specifically, Defendants’ Depo-Provera label in the EU now contains the following addition
15 under the section titled “**Special warnings and precautions for use**”: “Meningioma: Meningiomas
16 have been reported following long term administration of progestogens, including
17 medroxyprogesterone acetate. Depo-Provera should be discontinued if a meningioma is diagnosed.
18 Caution is advised when recommending Depo-Provera to patients with a history of meningioma.”

20 117. Additionally, Defendants’ Package Leaflet in the EU which provides information for
21 the patient states that “before using Depo-Provera[,]... it is important to tell your doctor or healthcare
22 professional if you have, or have ever had in the past ... a meningioma (a usually benign tumor that
23 forms in the layers of tissue that cover your brain and spinal cord).”
24

26 ²⁸ “Hormone medication could increase risk of brain tumours, French study finds,” The Guardian,
27 published online Mar. 27, 2024 (available at
28 <https://www.theguardian.com/society/2024/mar/27/hormone-medication-brain-tumours-risk-progestogens-study>) (last accessed Sept. 12, 2024).

1 118. Nothing was or is stopping Defendants from adding similar language to the label and
2 package insert for Depo-Provera in the United States. Defendants could have at any time made
3 “moderate changes” to the label.

4 119. Specifically, Defendants could have filed a “Changes Being Effected” (“CBE”)
5 supplement under Section 314.70(c) of the FDCA to make “moderate changes” to Depo-Provera’s
6 label without any prior FDA approval.

7 120. Examples of moderate label changes that can be made via a CBE supplement explicitly
8 include changes “to reflect newly acquired information” in order to “add or strengthen a
9 contraindication, warning, precaution, or adverse reaction.” By definition and by regulation such
10 changes to add a warning based on newly acquired information—such as that imparted by newly
11 emerging literature like the litany of studies cited above—are considered a “moderate change.” §
12 340.70(c)(6)(iii).
13

14 121. Recently, the Third Circuit reaffirmed that plain text interpretation of the CBE
15 supplement process in a precedential decision holding that the defendant in that case, Merck, could not
16 rely on a preemption defense based on an allegedly irreconcilable conflict between federal (FDCA)
17 and state (civil tort) law so long as the warning could have been effected via a CBE change. *See*
18 *generally In re Fosamax (Alendronate Sodium) Prods. Liab. Litig.*, Case No. 22-3412, D.I. 82 at 73 on
19 the docket (J. Jordan) (3d Cir. Sept. 20, 2024) (noting “the availability of a label change via a CBE
20 supplement is problematic for Merck, as will very often be the case for pharmaceutical companies
21 raising an impossibility defense”).
22

23 122. Defendants could have also instructed physicians to consider its own safer alternative
24 design, a lower dose medroxyprogesterone acetate injected subcutaneously instead of the more
25 invasive and painful intramuscular injection method. Studies going back at least ten years have shown
26 that the 150 mg dose of Depo-Provera—when administered subcutaneously, instead of
27
28

1 intramuscularly—is absorbed by the body at a similarly slower rate as the lower dose 104 mg Depo-
2 SubQ Provera 104 version and never exceeds more than a small fraction of the dangerously high serum
3 levels seen in the first several days with intramuscular administration of 150 mg Depo-Provera.²⁹
4 Nevertheless, Defendants never produced a 150 mg subcutaneous version.

5 123. Another study published in *Contraception: X* in 2022 concluded that not only was the
6 lower dose Depo-SubQ Provera 104 just as effective as 150 mg Depo-Provera when administered
7 properly, but it could also be administered every 16 weeks instead of every 12 weeks due to the more
8 gradual uptake of the subcutaneous administration route. That same study found that 150 mg Depo-
9 Provera if injected subcutaneously could remain at efficacious levels in the blood for even longer, up
10 to six (6) months.³⁰

12 124. As with subcutaneously administered Depo-SubQ Provera 104, the study authors noted
13 “subcutaneous administration of 150 mg Depo-Provera every 6 months would be a highly effective
14 repurposing ... with a similar reduction in cumulative exposure.” The authors concluded: “The use of
15 an unnecessarily high exposure to limit the residual chance of treatment failure would be a disservice
16 to the vast majority of women if a lower exposure can reduce side effects, costs, or otherwise make the
17 product more acceptable.”³¹

19 125. Despite knowing the subcutaneous administration of 150 mg Depo-Provera would have
20 resulted in much less risk of dangerous side effects like meningioma while providing the same
21 contraceptive efficacy for twice as long (and therefore would have required only half as many doses of
22 Defendants’ product per year), Defendants failed to produce a 150 mg subcutaneous version.

25 ²⁹ See Shelton, et al., “Subcutaneous DPMA: a better low dose approach,” *Contraception*, Vol. 89, pp.
26 341-43 (2014).

27 ³⁰ See Taylor, et al., “Ovulation suppression following subcutaneous administration of depot
medroxyprogesterone acetate,” *Contraception: X*, Vol. 4 (2022).

28 ³¹ *Id.*

1 126. Knowing that the lower dose 104 mg Depo-SubQ Provera 104 was equally effective
2 and was easier to administer since it involved a smaller needle being injected only below the skin and
3 not all the way into the muscle, Defendants could have educated the gynecology community that it
4 already had a safer alternative product to 150 mg Depo-Provera, which was more well known to
5 prescribers and patients.

6 127. In Europe and other countries outside of the United States, this 104 mg subcutaneous
7 dose has a more accessible trade name, “Sayana Press”, unlike the unwieldy proprietary developmental
8 name of “Depo-SubQ Provera 104”. Sayana Press as sold in Europe may be self-administered by
9 patients, obviating the need for quarterly visits to a medical practitioner.

11 128. When Depo-SubQ Provera 104, under NDA number 21-583, submitted by Defendant
12 Pharmacia & Upjohn, a subsidiary of Defendant Pfizer, was approved by the FDA on February 17,
13 2004, more than two decades ago, those Defendants submitted a proposed trade name that the FDA did
14 not approve, so instead, the proprietary name Depo-SubQ Provera 104 was deemed to be the brand
15 name.

16 129. Inexplicably, and presumably for commercially beneficial or contractual reasons,
17 Defendant Pfizer made a conscious decision to not seek an alternative commercially more accessible
18 brand name, and to not endeavor to more vigorously advocate for the sale of Depo-SubQ Provera 104
19 to patients seeking contraception, despite knowing it had a lower safer and effective dosage which
20 would mitigate the potential for adverse reactions engendered by a high dose progestin, including the
21 risk of developing or worsening meningioma tumors.

22 130. The “lowest effective dose” is a well-known concept in the field of pharmaceuticals
23 wherein a drug-maker should seek to find the lowest possible dose at which the drug of interest is
24 efficacious for the intended use, as any additional dosage on top of that lowest effective dose is
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1 inherently superfluous and can only increase the risk of unwanted and potentially dangerous side
2 effects while providing no additional efficacy.

3 131. Either change—adding a warning about the risk of meningioma based on “newly
4 acquired information,” or, advising physicians to consider a switch to subcutaneous Depo-SubQ
5 Provera 104—either on its own, or taken together, would have constituted a “moderate change”
6 justifying a simple CBE supplement that Defendants could have effectuated immediately and simply
7 notified the FDA thereafter. Yet, Defendants have failed to do so, and that failure continues to date.
8

9 132. Defendants ignored reports from patients and health care providers throughout the
10 United States which indicated that Depo-Provera failed to perform as intended. Defendants also
11 knew or should have known of the effects associated with long term use of Depo-Provera, which led
12 to the severe and debilitating injuries suffered by Plaintiff and numerous other patients. Rather
13 than conducting adequate testing to determine the cause of these injuries for which it had notice or
14 rule out Depo-Provera’s design as the cause of the injuries, Defendants continued to falsely and
15 misleadingly market Depo-Provera as a safe and effective prescription drug for contraception and
16 other indications.
17

18 133. Defendants’ Depo-Provera was at all times utilized and prescribed in a manner
19 foreseeable to Defendants, as Defendants generated the instructions for use for Plaintiff to receive
20 Depo-Provera injections.

21 134. Plaintiff and Plaintiff’s physicians foreseeably used Depo-Provera, and did not
22 misuse or alter Depo-Provera in an unforeseeable manner.
23

24 135. Through its affirmative misrepresentations and omissions, Defendants actively
25 concealed from Plaintiff and her physicians the true and significant risks associated with Depo-
26 Provera use.
27
28

1 136. As a result of Defendants' actions, Plaintiff and her physicians were unaware, and
2 could not have reasonably known or have learned through reasonable diligence, that Plaintiff would
3 be exposed to the risks identified in this Complaint and that those risks were the direct and
4 proximate result of Defendants' conduct.

5 137. As a direct result of being prescribed and consuming Depo-Provera, Plaintiff has
6 been permanently and severely injured, having suffered serious consequences.

7 138. As a direct and proximate result of her Depo-Provera use, Plaintiff suffered severe
8 mental and physical pain and suffering and has sustained permanent injuries and emotional distress,
9 along with economic loss including past and future medical expenses.

10 139. Despite diligent investigation by Plaintiff into the cause of these injuries, including
11 consultations with medical providers, the nature of Plaintiff's injuries and damages and their
12 relationship to Depo-Provera was not discovered, and through reasonable care and diligence could not
13 have been discovered, until a date within the applicable statute of limitations for filing Plaintiff's
14 claims.
15

16
17 **LIABILITY OF PFIZER, GREENSTONE, VIATRIS, AND PRASCO FOR THE**
18 **"AUTHORIZED GENERICS"**

19 140. Defendants Greenstone, Viatris, and Prasco were at different times from 2004 until the
20 present the authorized generic "manufacturer" and distributor operating under the same NDA of Depo-
21 Provera, with the express permission of Pfizer, to make, label, distribute, sell, and market Depo-
22 Provera without the brand name on its label, even though it is the exact same drug product as the
23 branded Depo-Provera manufactured in some or all instances by Pfizer.

24 141. Accordingly, the authorized generic distributors Greenstone, Viatris, and Prasco
25 operated as if they were the brand name holder under the same NDA and could have changed the brand
26 name label to warn of the risks of meningioma and the use of high dose progestins.
27
28

1 142. Further, the “authorized generics” distributors Greenstone, Viatris, and Prasco could
2 have requested that Pfizer, with whom they were under contract to sell the “authorized generic”, to
3 change the brand name label to warn of the risks of meningioma and the use of high dose progestins.

4 143. Pfizer had a duty to change the label knowing that its “authorized generic” distributors
5 Greenstone, Viatris, and Prasco, with whom they were in contract and receiving revenue from the sale
6 of the “authorized generic” DMPA, were selling the “authorized generic” without warning of
7 meningioma risk.
8

9 144. Pfizer knew that its authorized generic manufacturers held a large market share of its
10 manufactured Depo-Provera under a different name.

11 145. Pfizer was at some or all of the pertinent times the actual manufacturer of the DMPA,
12 identical to Depo-Provera other than its name, which was sold by Defendants Greenstone, Viatris, and
13 Prasco who were at different times the “authorized generic” distributor, with the express permission
14 of Pfizer, to distribute, sell, and market Depo-Provera without the brand name on its label.
15

16 **INNOVATOR LIABILITY UNDER CALIFORNIA LAW**
17

18 146. In October of 2002, Defendant Pfizer's patent for Depo-Provera expired. Following
19 this, the FDA approved various generic versions of Depo-Provera for sale in the United States. Despite
20 the availability of generics, Pfizer has continued to manufacture, market, and distribute the brand-
21 name Depo-Provera across the United States, including in California.

22 147. A manufacturer wishing to market a generic version of an FDA-approved drug can
23 submit an Abbreviated New Drug Application (ANDA). This allows the generic manufacturer to rely
24 on the NDA filed by the brand-name manufacturer by demonstrating that the generic version contains
25 the same active ingredients and is biologically equivalent to the brand-name drug.³²
26

27 _____
28 ³² See 21 U.S.C. § 355(j)(2)(A)(ii), (iv).

1 148. As part of the NDA, the brand-name manufacturer must propose the exact text of the
2 label, subject to FDA approval.³³ For generics, the ANDA process mandates that the safety and
3 efficacy labeling must be identical to that of the brand-name drug.³⁴

4 149. While the brand-name manufacturer bears responsibility for the accuracy and adequacy
5 of the drug label, generic manufacturers are only required to ensure that their labels mirror the brand-
6 name version.³⁵ The California Supreme Court has reasoned that because a brand-name manufacturer
7 is responsible for the content of a drug's warning label, it “knows to a legal certainty ... that any
8 deficiencies in the label for its drug will be perpetrated in the label for its generic bioequivalent.”³⁶ As
9 a result, the content of the generic labels for Depo-Provera bioequivalents is entirely dictated by the
10 brand-name manufacturer Defendant Pfizer’s label. Thus, California law liability for failure to warn
11 can extend to Defendant Pfizer, even when the consumer is prescribed only the generic version.
12

13 150. Because generic manufacturers must replicate the brand-name label exactly, Defendant
14 Pfizer exerted exclusive control over the contents of the labels used by generic versions of Depo-
15 Provera that Plaintiff may have been prescribed and administered. Consequently, any deficiencies or
16 omissions in Defendant Pfizer’s label would have been reflected in the generic labels.
17

18 151. As the brand-name manufacturer of Depo-Provera, Defendant Pfizer had and continues
19 to have a duty to ensure that the labeling for Depo-Provera remains accurate and adequate “as soon as
20 there is reasonable evidence of an association of a serious hazard with a drug,” regardless of whether
21 a causal relationship has been established.³⁷ Defendant Pfizer was not only in the best position to
22 provide warnings regarding Depo-Provera's risks but was also the only entity legally authorized to
23 update the label unilaterally under federal law.
24

25 _____
³³ See 21 U.S.C. § 355; see also 21 C.F.R. § 314.105(b).

26 ³⁴ See 21 U.S.C.A. § 355(j); see also *PLIVA, Inc. v. Mensing*, 564 U.S. 604, 612-13 (2011).

27 ³⁵ See generally 21 U.S.C. § 355; see also 21 C.F.R. § 314.105(b).

28 ³⁶ *T.H. v. Novartis Pharm. Corp.*, 4 Cal. 5th 145, at 166 (2017).

³⁷ See 21 C.F.R. § 201.80(e).

1 152. Defendant Pfizer knew or should have known that any failure to adequately warn of
2 Depo-Provera's risks would be replicated in the labels of its generic bioequivalents, directly affecting
3 the information available to physicians and patients regarding both the brand-name and generic drugs.
4 Accordingly, it is foreseeable that the warnings included or omitted on the brand-name drug label
5 would influence dispensing of the generic drug and the decision-making of unsuspecting doctors and
6 patients, like Plaintiff and Plaintiff's physicians, as to whether to take a generic equivalent of Depo-
7 Provera and/or brand-named Depo-Provera for contraception.

8
9 153. As the brand-name manufacturer of Depo-Provera, Defendant Pfizer could have, at any
10 time, unilaterally updated the Depo-Provera label without waiting for FDA preapproval in order to
11 "add or strengthen a contraindication, warning, precaution, or adverse reaction" under the CBE
12 regulation.³⁸ As the brand name manufacturer of Depo-Provera, Defendant Pfizer had a duty to give
13 information about Depo-Provera to the medical community and public at large.

14 154. Despite having the ability and obligation to provide timely and adequate warnings,
15 Defendant Pfizer failed to take such action, contributing to the harm suffered by Plaintiff.

16
17 155. Thus, to the extent that any of the approximately sixty-four (64) doses of Depo-Provera
18 administered to Plaintiff were generic, Defendant Pfizer is additionally liable for any resultant harm
19 to Plaintiff from those generic doses under California's well-established doctrine of innovator liability.

20 **EQUITABLE TOLLING OF STATUTE OF LIMITATIONS**

21
22 156. Defendants willfully, wantonly, and intentionally conspired, and acted in concert, to
23 withhold information from Plaintiff, Plaintiff's healthcare providers, and the general public concerning
24 the known hazards associated with the use of, and exposure to, Depo-Provera, particularly over
25 extended periods of time.

26
27
28 ³⁸ See 21 C.F.R. § 314.70(c)(6)(iii)(A).

1 157. Defendants willfully, wantonly, and intentionally conspired, and acted in concert, to
2 withhold safety-related warnings from the Plaintiff, and the general public concerning the known
3 hazards associated with the use of, and exposure to, Depo-Provera, particularly over extended periods
4 of time.

5 158. Defendants willfully, wantonly, and intentionally conspired, and acted in concert, to
6 withhold instructions from the Plaintiff, her family members, and the general public concerning how
7 to identify, mitigate, and/or treat known hazards associated with the use of, and exposure to, Depo-
8 Provera, particularly over extended periods of time.

9 159. The aforementioned studies reveal that discontinuing use of high dose progesterone
10 and progestin, including Depo-Provera, can retard the growth of meningiomas, but failed to warn the
11 medical community and the Plaintiff of this method to mitigate the damage of a developing
12 meningioma.

13 160. Defendants willfully, wantonly, and intentionally conspired, and acted in concert, to
14 ignore relevant safety concerns and to deliberately not study the long-term safety and efficacy of Depo-
15 Provera, particularly in chronic long-term users of Depo-Provera.

16 161. Defendants failed to disclose a known defect and, instead, affirmatively misrepresented
17 that Depo-Provera was safe for its intended use. Defendants disseminated labeling, marketing,
18 promotion and/or sales information to Plaintiff, her healthcare providers, and the general public
19 regarding the safety of Depo-Provera knowing such information was false, misleading, and/or
20 inadequate to warn of the safety risks associated with long-term Depo-Provera use. Defendants did so
21 willfully, wantonly, and with the intent to prevent the dissemination of information known to them
22 concerning Depo-Provera's safety.

23 162. Further, Defendants actively concealed the true risks associated with the use of Depo-
24 Provera, particularly as they relate to the risk of serious intracranial meningioma, by affirmatively
25

1 representing in numerous communications, which were disseminated to Plaintiff, her healthcare
2 providers, and which included, without limitation, the Package Insert and the Medication Guide, that
3 there were no warnings required to safely prescribe and take Depo-Provera and no intracranial
4 meningioma-related adverse side effects associated with use of Depo-Provera.

5 163. Due to the absence of any warning by the Defendants as to the significant health and
6 safety risks posed by Depo-Provera, Plaintiff was unaware that Depo-Provera could cause the
7 development of a serious and debilitating intracranial meningioma, as this danger was not known to
8 Plaintiff, Plaintiff's healthcare providers, or the general public.

9 164. Due to the absence of any instructions for how to identify and/or monitor Depo-Provera
10 patients for potential intracranial meningioma-related complications, Plaintiff was unaware that Depo-
11 Provera could cause serious, intracranial meningioma-related injuries, as this danger was not known
12 to Plaintiff, Plaintiff's healthcare providers, or the general public.

13 165. Given Defendants' conduct and deliberate actions designed to deceive Plaintiff,
14 Plaintiff's healthcare providers, and the general public, with respect to the safety and efficacy of Depo-
15 Provera, Defendants are estopped from relying on any statute of limitations defenses.

16
17
18
19 **CONDUCT WARRANTING PUNITIVE DAMAGES**

20 166. For the reasons set forth above and addressed below, Defendant Pfizer acted with a
21 conscious disregard of the safety of Plaintiff and all the other women, many who were young and of
22 lower socioeconomic status, who were subjected to high dose injections of 150 mg Depo-Provera with
23 the known and/or knowable risk of meningioma brain tumors which was generally accepted in the
24 scientific community, while Defendant Pfizer had available its very own safer alternative medication,
25 Depo Sub-Q Provera 104. Exemplary damages are warranted to punish and deter Defendant Pfizer
26 and others from such conduct in the future.
27
28

COUNT I

STRICT LIABILITY – FAILURE TO WARN

167. Plaintiff incorporates by reference each and every preceding paragraph as though fully set forth herein.

168. At all times material herein, Defendants engaged in the business of researching, testing, developing, manufacturing, labeling, marketing, selling, inspecting, handling, storing, distributing, and/or promoting Depo-Provera and placed Depo-Provera into the stream of commerce in a defective and unreasonably dangerous condition. These actions were under the ultimate control and supervision of Defendants.

169. Defendants, as manufacturers, distributors, and marketers of pharmaceutical drugs, are held to the level of knowledge of an expert in the field, and further, Defendants knew or should have known based on information that was available and generally accepted in the scientific community that warnings and other clinically relevant information and data which they distributed regarding the risks associated with the use of Depo-Provera were inadequate.

170. Plaintiff and Plaintiff's treating physicians did not have the same knowledge as Defendants and no adequate warning or other clinically relevant information or data was communicated to Plaintiff or to Plaintiff's treating physicians.

171. Defendants had and continue to have a duty to provide adequate warnings and instructions for Depo-Provera, to use reasonable care to design a product that is not unreasonably dangerous to users, and to adequately understand, test, and monitor their product.

172. Defendants had and continue to have a duty to provide consumers, including Plaintiff and Plaintiff's physicians, with warnings and other clinically relevant information and data generally accepted within the scientific community regarding the risks and dangers associated with Depo-Provera, as it became or could have become available to Defendants.

1 173. Defendants marketed, promoted, distributed and sold an unreasonably dangerous and
2 defective prescription drug, Depo-Provera, to health care providers empowered to prescribe and
3 dispense Depo-Provera, to consumers, including Plaintiff, without adequate warnings and other
4 clinically relevant information and data regarding the risk of meningioma and the risks of
5 unnecessarily excessive progestin exposure which was available and generally accepted within the
6 scientific community. Through both omission and affirmative misstatements, Defendants misled the
7 medical community about the risk and benefit balance of Depo-Provera, which resulted in injury to
8 Plaintiff.
9

10 174. Defendants knew or should have known through testing, scientific knowledge,
11 advances in the field, published research in major peer-reviewed journals, or otherwise, that Depo-
12 Provera created a risk of developing serious and debilitating intracranial meningioma. At all relevant
13 times this information was readily available and generally accepted within the scientific community.
14

15 175. Despite the fact that Defendants knew or should have known based on information
16 generally accepted within the scientific community that Depo-Provera with its higher than needed
17 progestin dosage caused unreasonable and dangerous side effects, they continue to promote and
18 market Depo-Provera without providing adequate clinically relevant information and data or
19 recommending patients be monitored.

20 176. Defendants knew that a safer alternative design and product existed, including its own
21 Depo-SubQ Provera 104 which contained substantially less progestin but was equally effective in
22 preventing pregnancy, but failed to warn the medical community and the patients about the risks of
23 the high dose which could be mitigated by using the lower dose formulation, Depo-SubQ Provera 104.
24

25 177. Defendants knew or should have known that consumers, and Plaintiff, specifically,
26 would foreseeably and needlessly suffer injury as a result of Defendants' failures.
27
28

1 178. The Depo-Provera supplied to Plaintiff by Defendants was defective, unreasonably
2 dangerous, and had inadequate warnings or instructions at the time it was sold, and Defendants also
3 acquired additional knowledge and information confirming the defective and unreasonably dangerous
4 nature of Depo-Provera. Despite this knowledge and information, Defendants failed and neglected to
5 issue adequate warnings that Depo-Provera causes serious and potentially debilitating intracranial
6 meningioma and/or instructions concerning the need for monitoring and potential discontinuation of
7 use of Depo-Provera.
8

9 179. Defendants' failure to provide adequate warnings or instructions rendered Depo-
10 Provera unreasonably dangerous in that it failed to perform as safely as an ordinary patient, prescriber,
11 and/or other consumer would expect when used as intended and/or in a manner reasonably foreseeable
12 by the Defendants, and in that the risk of danger outweighs the benefits.

13 180. Defendants failed to provide timely and adequate warnings to physicians, pharmacies,
14 and consumers, including Plaintiff and Plaintiff's intermediary physicians.
15

16 181. Plaintiff's various prescribing physicians, nurse practitioners, physician assistants, and
17 nurses (hereinafter collectively referred to as "Plaintiff's Prescribing and Administering Health Care
18 Providers") would not have prescribed and administered Depo-Provera to Plaintiff had they been
19 apprised by Defendants of the unreasonably high risk of meningioma associated with usage of Depo-
20 Provera.

21 182. Alternatively, even if Defendants had apprised Plaintiff's Prescribing and
22 Administering Health Care Providers of the unreasonably high risk of meningioma associated with
23 usage of Depo-Provera and these Prescribing and Administering Health Care Providers had still
24 recommended usage of Depo-Provera to Plaintiff, the Prescribing and Administering Health Care
25 Providers would have relayed the information concerning the risk of meningioma to Plaintiff, and the
26 alternative treatment of the lower dose subcutaneous Depo-SubQ Provera 104, and Plaintiff as an
27
28

1 objectively prudent person would not have chosen to take Depo-Provera, and/or would have opted to
2 take safer and lower dose Depo-SubQ Provera 104, notwithstanding Plaintiff's Prescribing Physician
3 and Administering Health Care Providers' continued recommendation.

4 183. Similarly, if Defendants had warned of the unreasonably high risk of meningioma
5 associated with the usage of Depo-Provera, and the availability of the safer and equally effective lower
6 dose Depo-SubQ Provera 104 in the Patient Information handout, Plaintiff as an objectively prudent
7 person would not have chosen to take Depo-Provera, and/or would have opted to take the safer, lower,
8 and equally effective dose of Depo-SubQ Provera 104, notwithstanding Plaintiff's Prescribing and
9 Administering Health Care Providers' recommendation.
10

11 184. Defendants failed to include adequate warnings and/or provide adequate clinically
12 relevant information and data that would alert Plaintiff and Plaintiff's Prescribing and Administering
13 Health Care Providers of the dangerous risks of Depo-Provera including, among other things, the
14 development of intracranial meningioma.
15

16 185. Defendants failed to provide adequate post-marketing warnings and instructions after
17 Defendants knew or should have known of the significant risks of, among other things, intracranial
18 meningioma.

19 186. Defendants continued to aggressively promote and sell Depo-Provera, even after they
20 knew or should have known of the unreasonable risks of intracranial meningioma caused by the drug.

21 187. Defendants had an obligation to provide Plaintiff and Plaintiff's Prescribing and
22 Administering Health Care Providers with adequate clinically relevant information and data and
23 warnings regarding the adverse health risks associated with exposure to Depo-Provera, and/or that
24 there existed safer and more or equally effective alternative drug products.
25

26 188. By failing to adequately test and research harms associated with Depo-Provera, and by
27 failing to provide appropriate warnings and instructions about Depo-Provera use, patients and the
28

1 medical community, including prescribing doctors, were inadequately informed about the true risk-
2 benefit profile of Depo-Provera and were not sufficiently aware that serious and potentially
3 debilitating intracranial meningioma might be associated with use of Depo-Provera. Nor were the
4 medical community, patients, patients' families, or regulators appropriately informed that serious and
5 potentially debilitating intracranial meningioma might be a side effect of Depo-Provera and should or
6 could be reported as an adverse event.

7
8 189. The Depo-Provera products designed, researched, manufactured, tested, advertised,
9 promoted, marketed, sold and distributed by Defendants were defective due to inadequate post-
10 marketing surveillance and/or warnings because, even after Defendants knew or should have known
11 of the risks of severe and permanent intracranial meningioma-related injuries from ingesting Depo-
12 Provera, Defendants failed to provide adequate warnings to users or consumers of the products, and
13 continued to improperly advertise, market and/or promote Depo-Provera.

14
15 190. Depo-Provera is defective and unreasonably dangerous to Plaintiff and other consumers
16 regardless of whether Defendants had exercised all possible care in its preparation and sale.

17
18 191. The foreseeable risk of serious and potentially debilitating intracranial meningioma
19 caused by Depo-Provera could have been reduced or avoided by Plaintiff, prescribers, and/or other
20 consumers had Defendants provided reasonable instructions or warnings of these foreseeable risks of
21 harm.

22
23 192. As a direct and proximate result of Defendants' conduct, including the inadequate
24 warnings, dilution or lack of information, lack of adequate testing and research, and the defective and
25 dangerous nature of Depo-Provera, Plaintiff suffered bodily injuries and resulting pain and suffering,
26 disability, mental anguish, loss of capacity for the enjoyment of life, expense of medical and nursing
27 care and treatment, loss of earnings, loss of ability to earn money and other economic losses, and
28

1 aggravation of previously existing conditions. The losses are either permanent or continuing, and
2 Plaintiff will suffer the losses in the future.

3
4 **COUNT II**

5 **STRICT LIABILITY – DESIGN DEFECT**

6 193. Plaintiff incorporates by reference each and every preceding paragraph as though fully
7 set forth herein.

8 194. At all times material herein, Defendants engaged in the business of researching, testing,
9 developing, manufacturing, labeling, marketing, selling, inspecting, handling, storing, distributing,
10 and/or promoting Depo-Provera and placed Depo-Provera into the stream of commerce in a defective
11 and unreasonably dangerous condition. These actions were under the ultimate control and supervision
12 of Defendants.

13 195. Defendants, as manufacturers, designers, distributors, and marketers of pharmaceutical
14 drugs, had a duty to design a product free from a defective condition that was unreasonably dangerous
15 to Plaintiff.

16 196. Depo-Provera was designed in such a way, using such a high dose of progesterone not
17 necessary for effective contraception, that it posed an unreasonable risk of intracranial meningioma
18 and by placing and keeping Depo-Provera on the market despite Depo-Provera being in a defective
19 condition.
20

21 197. Depo-SubQ Provera 104 is a lower dosage version of Depo-Provera that contains 104
22 mg / 0.65mL and is injected subcutaneously every three (3) months. According to the label, Depo-
23 SubQ Provera 104 can be used for both contraception and treatment of endometriosis.

24 198. Depo-SubQ Provera 104 never attained meaningful market share, and Defendant failed
25 to promote the product to the medical community as a safer and equally effective method of
26 contraception for women choosing to receive quarterly injections.
27
28

1 199. Defendant failed to promote and encourage conversion of the prescribing
2 gynecological community to Depo-SubQ Provera 104, fearing that doing so could instill a concern of
3 safety as to the risks of its high dose progesterone long standing product, Depo-Provera.

4 200. It has long been a tenet in the medical and toxicological community that the “dose
5 makes the poison.” Defendants had a viable safer and lower dose alternative in Depo-SubQ Provera
6 104 but failed to warn the medical community prescribing and administering Depo-Provera that Depo-
7 SubQ Provera 104 was a safer alternative.

8 201. Moreover, the 150 mg Depo-Provera itself could have been a viable lower effective
9 dose if it had simply been designed, approved, and sold to be administered subcutaneously, like Depo-
10 SubQ Provera 104 is administered, instead of intramuscularly.

11 202. Injections given intramuscularly are well-known to be absorbed by the body and taken
12 up in the blood serum at much faster rates than injections given subcutaneously because of the much
13 higher vascularization of deep muscle tissue compared to the dermis.

14 203. Studies have shown that 150 mg Depo-Provera administered intramuscularly causes a
15 spike in blood serum levels of DMPA that is more than four (4) times higher than the peak blood
16 serum concentration of DMPA when that same 150 mg Depo-Provera shot is given subcutaneously,
17 and that very high intramuscular peak concentration persists for several days.³⁹ In fact, 150 mg Depo-
18 Provera administered subcutaneously has a remarkably similar pharmacokinetic profile to Depo-SubQ
19 Provera 104.⁴⁰

20 204. Thus, there are two lower effective doses of Depo-Provera—both Depo-SubQ Provera
21 104, *and* the very same 150 mg Depo-Provera simply given subcutaneously instead of intramuscularly.

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25
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27 ³⁹ See Shelton, et al., “Subcutaneous DPMA: a better low dose approach,” *Contraception*, Vol. 89, pp.
28 341-43 (2014).

⁴⁰ See *id.* at 342.

1 205. Defendants wantonly and willfully failed to apprise the public, including the FDA, the
2 medical community, Plaintiff, Planned Parenthood, and Plaintiff's physicians, of the greatly reduced
3 risk of meningioma when injecting 150 mg Depo-Provera subcutaneously compared to the indicated
4 method of intramuscular injection because Defendants did not want to raise any alarms with respect
5 to the safety profile of Depo-Provera and did not want to lose any of its lucrative market share held in
6 part through its contracts with "authorized generic" partners and subsidiaries.

7
8 206. Defendants knew or should have known that the Depo-Provera they developed,
9 manufactured, labeled, marketed, sold, and/or promoted was defectively designed in that it posed a
10 serious risk of severe and permanent intracranial-meningioma-related injuries when injected
11 intramuscularly.

12 207. Defendants have a continuing duty to design a product that is not unreasonably dangerous
13 to users and to adequately understand, test, and monitor their product.

14 208. Defendants sold, marketed and distributed a product that is unreasonably dangerous for
15 its normal, intended, and foreseeable use.

16
17 209. Defendants designed, researched, manufactured, tested, advertised, promoted,
18 marketed, sold and distributed Depo-Provera, a defective product which created an unreasonable risk
19 to the health of consumers, and Defendants are therefore strictly liable for the injuries sustained by
20 Plaintiff.

21 210. The Depo-Provera supplied to Plaintiff by Defendants was defective in design or
22 formulation in that, when it left the hands of the manufacturer or supplier, it was in an unreasonably
23 dangerous and a defective condition because it failed to perform as safely as an ordinary consumer
24 would expect when used as intended or in a manner reasonably foreseeable to Defendants, posing a
25 risk of serious and potentially debilitating intracranial meningioma to Plaintiff and other consumers.
26
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1 211. The Depo-Provera ingested by Plaintiff was expected to, and did, reach Plaintiff
2 without substantial change in the condition in which it is sold.

3 212. The Depo-Provera ingested by Plaintiff was in a condition not contemplated by the
4 Plaintiff in that it was unreasonably dangerous, posing a serious risk of permanent vision and retinal
5 injuries.

6 213. Depo-Provera is a medication prescribed for contraception and treatment of
7 endometriosis, among other uses. Depo-Provera in fact causes serious and potentially debilitating
8 intracranial meningioma, a brain tumor that can cause severe damage and require invasive surgical
9 removal, harming Plaintiff and other consumers.

11 214. Plaintiff, ordinary consumers, and prescribers would not expect a contraceptive drug
12 designed, marketed, and labeled for contraception to cause intracranial meningioma.

13 215. The Depo-Provera supplied to Plaintiff by Defendants was defective in design or
14 formulation in that, when it left the hands of the manufacturer or supplier, it had not been adequately
15 tested, was in an unreasonably dangerous and defective condition, provided an excessive dose of
16 progestin for its purpose and posed a risk of serious and potentially debilitating intracranial
17 meningioma to Plaintiff and other consumers.

19 216. The Depo-Provera supplied to Plaintiff by Defendants was defective in design or
20 formulation in that its effectiveness as a contraceptive did not outweigh the risks of serious and
21 potentially debilitating intracranial meningioma posed by the drug. In light of the utility of the drug
22 and the risk involved in its use, the design of the Depo-Provera drug makes the product unreasonably
23 dangerous.

25 217. Depo-Provera's design is more dangerous than a reasonably prudent consumer would
26 expect when used in its intended or reasonably foreseeable manner. It was more dangerous than
27 Plaintiff expected.

1 218. The intended or actual utility of Depo-Provera is not of such benefits to justify the risk
2 of intracranial meningioma which may cause severe and permanent injuries, thereby rendering the
3 product unreasonably dangerous.

4 219. The design defects render Depo-Provera more dangerous than other drugs and therapies
5 designed for contraception and causes an unreasonable increased risk of injury, including, but not
6 limited, to potentially debilitating intracranial meningioma and sequelae related thereto.

7 220. Defendants knew or should have known through testing, generally accepted scientific
8 knowledge, advances in the field, published research in major peer-reviewed journals, or other means,
9 that Depo-Provera created a risk of serious and potentially debilitating intracranial meningioma and
10 sequelae related thereto.

11 221. Depo-Provera is defective and unreasonably dangerous to Plaintiff and other
12 consumers in that, despite early indications and concerns that Depo-Provera use could result in vision
13 issues, Defendants failed to adequately test or study the drug, including but not limited to:
14 pharmacokinetics and pharmacodynamics of the drug, its effects on the development of brain tumors
15 like intracranial meningioma, the potential effects and risks of long-term use, the potential for inter-
16 patient variability, and/or the potential for a safer effective dosing regimen.

17 222. Defendants knew or should have known that consumers, Plaintiff specifically, would
18 foreseeably and needlessly suffer injury as a result of Depo-Provera's defective design.

19 223. Depo-Provera is defective and unreasonably dangerous to Plaintiff and other
20 consumers even if Defendants had exercised all possible care in the preparation and sale of Depo-
21 Provera.

22 224. As a direct and proximate result of Defendants' conduct and defective design, including
23 inadequate testing and research, and the defective and dangerous nature of Depo-Provera, Plaintiff
24 suffered bodily injuries that resulted in pain and suffering, disability, mental anguish, loss of capacity
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1 for the enjoyment of life, expense of medical and nursing care and treatment, loss of earnings, loss of
2 ability to earn money, and other economic losses. The losses are either permanent or continuing, and
3 Plaintiff will suffer losses in the future.

4 **COUNT III**

5 **NEGLIGENCE**

6
7 225. Plaintiff incorporates by reference each and every preceding paragraph as though fully
8 set forth herein.

9 226. At all times relevant herein, it was the duty of Defendants to use reasonable care in
10 the design, labeling, manufacturing, testing, marketing, distribution and/or sale of Depo-Provera.

11 227. Defendants failed to exercise ordinary care in the labeling, design, manufacturing,
12 testing, marketing, distribution and/or sale of Depo-Provera in that Defendants knew or should have
13 known that Depo-Provera created a high risk of unreasonable harm to Plaintiff and other users.

14 228. Defendants breached its duty of care to the Plaintiff and her physicians, in the testing,
15 monitoring, and pharmacovigilance of Depo-Provera.

16 229. In disregard of its duty, Defendants committed one or more of the following negligent
17 acts or omissions:
18

19
20 a. Manufacturing, producing, promoting, formulating, creating, developing,
21 designing, selling, and distributing Depo-Provera without thorough and adequate pre- and post-
22 market testing of the product;

23 b. Manufacturing, producing, promoting, advertising, formulating, creating,
24 developing, and designing, and distributing Depo-Provera while negligently and intentionally
25 concealing and failing to disclose clinical data which demonstrated the risk of serious harm
26 associated with the use of Depo-Provera;
27
28

1 c. Failing to undertake sufficient studies and conduct necessary tests to
2 determine whether or not Depo-Provera was safe for its intended use;

3 d. Failing to disclose and warn of the product defect to the regulatory agencies,
4 the medical community, and consumers that Defendants knew and had reason to know that Depo-
5 Provera was indeed unreasonably unsafe and unfit for use by reason of the product's defect and
6 risk of harm to its users;

7 e. Failing to warn Plaintiff, the medical and healthcare community, and
8 consumers of the known and knowable product's risk of harm which was unreasonable and that
9 there were safer and effective alternative products available to Plaintiff and other consumers;

10 f. Failing to provide adequate instructions, guidelines, and safety precautions to
11 those persons to whom it was reasonably foreseeable would use Depo-Provera;

12 g. Advertising, marketing, and recommending the use of Depo-Provera, while
13 concealing and failing to disclose or warn of the dangers known and knowable by Defendants to be
14 connected with, and inherent in, the use of Depo-Provera;

15 h. Representing that Depo-Provera was safe for its intended use when in fact
16 Defendants knew and should have known the product was not safe for its intended purpose;

17 i. Continuing to manufacture and sell Depo-Provera with the knowledge that
18 Depo-Provera was unreasonably unsafe and dangerous;

19 j. Failing to use reasonable and prudent care in the design, research,
20 testing, manufacture, and development of Depo-Provera so as to avoid the risk of serious harm
21 associated with the use of Depo-Provera;

22 k. Failing to design and manufacture Depo-Provera so as to ensure the
23 drug was at least as safe and effective as other similar products;

24 l. Failing to ensure the product was accompanied by proper and accurate
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28

1 warnings about monitoring for potential symptoms related to intracranial meningioma associated with
2 the use of Depo-Provera;

3 m. Failing to ensure the product was accompanied by proper and accurate
4 warnings about known and knowable adverse side effects associated with the use of Depo-Provera
5 and that use of Depo-Provera created a high risk of severe injuries; and

6 n. Failing to conduct adequate testing, including pre-clinical and clinical
7 testing, and post-marketing surveillance to determine the safety of Depo-Provera.

8 o. Failing to sell a product with the lowest effective dose knowing that there
9 were safer lower effective dose formulations.
10

11 230. A reasonable manufacturer, designer, distributor, promoter, or seller under the same or
12 similar circumstances would not have engaged in the aforementioned acts and omissions.

13
14 231. As a direct and proximate result of the Defendants' negligent testing, monitoring, and
15 pharmacovigilance of Depo-Provera, Defendants introduced a product that they knew or should have
16 known would cause serious and permanent injuries related to the development of intracranial
17 meningioma, and Plaintiff has been injured tragically and sustained severe and permanent pain,
18 suffering, disability, and impairment, loss of enjoyment of life, loss of care, comfort, and economic
19 damages.
20

21 232. As a direct and proximate result of one or more of the above-stated negligent acts by
22 Defendants, Plaintiff suffered bodily injuries and resulting pain and suffering, disability, mental
23 anguish, loss of capacity for the enjoyment of life, expense of medical and nursing care and treatment,
24 loss of earnings, loss of consortium, loss of ability to earn money and other economic losses. The
25 losses are either permanent or continuing, and Plaintiff will suffer losses in the future.
26

27 **COUNT IV**
28

NEGLIGENT FAILURE TO WARN

233. Plaintiff incorporates by reference each and every preceding paragraph as though fully set forth herein.

234. At all times material herein, Defendants had a duty to exercise reasonable care and had the duty of an expert in all aspects of the warning and post-sale warning to assure the safety of Depo-Provera when used as intended or in a way that Defendants could reasonably have anticipated, and to assure that the consuming public, including Plaintiff and Plaintiff's physicians, obtained accurate information and adequate instructions for the safe use or non-use of Depo-Provera.

235. Defendants' duty of care was that a reasonably careful designer, manufacturer, seller, importer, distributor and/or supplier would use under like circumstances.

236. Defendants had a duty to warn Plaintiff, Plaintiff's physicians, and consumers of Depo-Provera's known and knowable dangers and serious side effects, including serious and potentially debilitating intracranial meningioma, as it was reasonably foreseeable to Defendants that Depo-Provera could cause such injuries.

237. At all times material herein, Defendants failed to exercise reasonable care and knew, or in the exercise of reasonable care should have known, that Depo-Provera had inadequate instructions and/or warnings.

238. Each of the following acts and omissions herein alleged was negligently and carelessly performed by Defendants, resulting in a breach of the duties set forth above. These acts and omissions include, but are not restricted to:

a. Failing to accompany their product with proper and adequate warnings, labeling, or instructions concerning the potentially dangerous, defective, unsafe, and deleterious propensity of Depo-Provera and of the risks associated with its use, including the severity and potentially irreversible nature of such adverse effects;

1 b. Disseminating information to Plaintiff and Plaintiff 's physicians that was
2 negligently and materially inaccurate, misleading, false, and unreasonably dangerous to patients
3 such as Plaintiff;

4 c. Failing to provide warnings or other information that accurately reflected the
5 symptoms, scope, and severity of the side effects and health risks;

6 d. Failing to adequately test and/or warn about the use of Depo-Provera,
7 including, without limitations, the possible adverse side effects and health risks caused by the use
8 of Depo-Provera;

9 e. Failure to adequately warn of the risks that Depo-Provera could cause the
10 development of intracranial meningioma and sequelae related thereto;

11 f. Failure to adequately warn of the risk of serious and potentially irreversible
12 injuries related to the development of intracranial meningioma, a brain tumor;

13 g. Failure to instruct patients, prescribers, and consumers of the need for al
14 monitoring when taking Depo-Provera for symptoms potentially related to the development of
15 intracranial meningioma;

16 h. Failure to instruct patients, prescribers, and consumers of the need to
17 discontinue Depo-Provera in the event of symptoms potentially related to the development of
18 intracranial meningioma;

19 i. Failing to provide instructions on ways to safely use Depo-Provera to avoid
20 injury, if any;

21 j. Failing to explain the mechanism, mode, and types of adverse events
22 associated with Depo-Provera;

23 k. Failing to provide adequate training or information to medical care providers
24 for appropriate use of Depo-Provera and patients taking Depo-Provera; and
25
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28

1 l. Representing to physicians, including but not limited to Plaintiff's
2 prescribing physicians, that this drug was safe and effective for use.

3 m. Failing to warn that there is a safer feasible alternative with a lower effective
4 dose of progestin.

5 n. Failing to warn that the 150 mg dosage of progestin injected intramuscularly
6 was an excessive and thus toxic dose capable of causing and or substantially contributing to the
7 development and growth of meningioma tumors.

8
9 239. Defendants knew or should have known of the risk and danger of serious bodily
10 harm from the use of Depo-Provera but failed to provide an adequate warning to patients and
11 prescribing physicians for the product, including Plaintiff and Plaintiff's prescribing physicians,
12 despite knowing the product could cause serious injury.

13 240. Plaintiff was prescribed and used Depo-Provera for its intended purpose.

14 241. Plaintiff could not have known about the dangers and hazards presented by Depo-
15 Provera.

16
17 242. The warnings given by Defendants were not accurate, clear, or complete and/or
18 were ambiguous.

19 243. The warnings, or lack thereof, that were given by Defendants failed to properly
20 warn prescribing physicians, including Plaintiff's prescribing physician, of the known and
21 knowable risk of serious and potentially irreversible injuries related to the development of
22 intracranial meningioma, and failed to instruct prescribing physicians to test and monitor for the
23 presence of the injuries and to discontinue use when symptoms of meningioma manifest.

24
25 244. The warnings that were given by the Defendants failed to properly warn Plaintiff
26 and prescribing physicians of the prevalence of intracranial meningioma and sequelae related
27 thereto.

1 250. At all times material herein, Defendants failed to exercise reasonable care and the duty
2 of an expert and knew, or in the exercise of reasonable care should have known, that Depo-Provera
3 was not properly manufactured, designed, compounded, tested, inspected, packaged, distributed,
4 marketed, advertised, formulated, promoted, examined, maintained, sold, prepared, or a combination
5 of these acts.

6 251. Each of the following acts and omissions herein alleged was negligently and carelessly
7 performed by Defendants, resulting in a breach of the duties set forth above. These acts and omissions
8 include, but are not restricted to negligently and carelessly:
9

10 a. Failing to use due care in developing, testing, designing, and manufacturing
11 Depo-Provera so as to avoid the aforementioned risks to individuals when Depo-Provera was being
12 used for contraception and other indications;
13

14 b. Failing to conduct adequate pre-clinical and clinical testing and post-
15 marketing surveillance to determine the safety of Depo-Provera; and
16

17 c. Designing, manufacturing, and placing into the stream of commerce a
18 product which was unreasonably dangerous for its reasonably foreseeable use, which Defendants
19 knew or should have known could cause injury to Plaintiff.
20

21 d. Failing to use due care in developing, testing, designing, and manufacturing
22 Depo-Provera with the lowest effective dose as a safer alternative which clearly existed at all
23 relevant times so as to avoid the aforementioned risks to individuals when high dose progestin
24 Depo-Provera was being used for contraception.

25 252. Defendants' negligence and Depo-Provera's failures arise under circumstances
26 precluding any other reasonable inference other than a defect in Depo-Provera.
27
28

1 the public and medical community, including Plaintiff and her Prescribing and Administering Health
2 Care Providers to request, recommend, purchase, and prescribe Depo-Provera.

3 259. The Defendants had a duty to accurately and truthfully represent to the medical and
4 healthcare community, medical device manufacturers, Plaintiff, her Prescribing and Administering
5 Health Care Providers and the public, the known risks of Depo-Provera, including its propensity to
6 cause intracranial meningioma and sequelae related thereto.
7

8 260. Defendants made continued omissions in the Depo-Provera labeling, including
9 promoting it as safe and effective while failing to warn of its propensity to cause intracranial
10 meningioma and sequelae related thereto.
11

12 261. Defendants made additional misrepresentations beyond the product labeling by
13 representing Depo-Provera as safe and effective for contraception and other indications with only
14 minimal risks.
15

16 262. Defendants misrepresented and overstated the benefits of Depo-Provera to Plaintiff,
17 Plaintiff's Prescribing and Administering Health Care Providers, and the medical community without
18 properly advising of the known risks associated with intracranial meningioma and sequelae related
19 thereto.

20 263. Defendants misrepresented and overstated that the Depo-Provera dosage was needed
21 to protect against pregnancy when Defendants knew that a safer alternative existed with forty-six (46)
22 fewer mg per dose of the powerful progestin being ingested quarterly in women, and when Defendants
23 could have warned and recommended usage of Depo-SubQ Provera 104 instead.
24

25 264. In reliance upon the false and negligent misrepresentations and omissions made by the
26 Defendants, Plaintiff and Plaintiff's Prescribing and Administering Health Care Providers were
27
28

1 induced to, and did use Depo-Provera, thereby causing Plaintiff to endure severe and permanent
2 injuries.

3 265. In reliance upon the false and negligent misrepresentations and omissions made by the
4 Defendants, Plaintiff and Plaintiff's Prescribing and Administering Health Care Providers were unable
5 to associate the injuries sustained by Plaintiff with her Depo-Provera use, and therefore unable to
6 provide adequate treatment. Defendants knew or should have known that the Plaintiff, Plaintiff's
7 Prescribing and Administering Health Care Providers, and the general medical community did not
8 have the ability to determine the true facts which were intentionally and/or negligently concealed and
9 misrepresented by the Defendants.
10

11 266. Plaintiff and her Prescribing and Administering Health Care Providers would not have
12 used or prescribed Depo-Provera had the true facts not been concealed by the Defendants.

13 267. Defendants had sole access to many of the material facts concerning the defective
14 nature of Depo-Provera and its propensity to cause serious and dangerous side effects.
15

16 268. At the time Plaintiff was prescribed and administered Depo-Provera, Plaintiff and her
17 Prescribing and Administering Health Care Providers were unaware of Defendants' negligent
18 misrepresentations and omissions.

19 269. The Defendants failed to exercise ordinary care in making representations concerning
20 Depo-Provera while they were involved in their manufacture, design, sale, testing, quality assurance,
21 quality control, promotion, marketing, labeling, and distribution in interstate commerce, because the
22 Defendants negligently misrepresented Depo-Provera's significant risk of unreasonable and
23 dangerous adverse side effects.
24

25 270. Plaintiff and Plaintiff's Prescribing and Administering Health Care Providers
26 reasonably relied upon the misrepresentations and omissions made by the Defendants, where the
27
28

1 concealed and misrepresented facts were critical to understanding the true dangers inherent in the use
2 of Depo-Provera.

3 271. Plaintiff and Plaintiff's Prescribing and Administering Health Care Providers' reliance
4 on the foregoing misrepresentations and omissions was the direct and proximate cause of Plaintiff's
5 injuries.

6 272. As a direct and proximate result of reliance upon Defendants' negligent
7 misrepresentations, Plaintiff suffered bodily injuries and resulting pain and suffering, disability,
8 mental anguish, loss of capacity for the enjoyment of life, expense of medical and nursing care and
9 treatment, loss of earnings, loss of consortium, loss of ability to earn money and other economic losses.
10 The losses are either permanent or continuing, and Plaintiff will suffer the losses in the future.
11

12 **COUNT VII**

13 **FRAUDULENT MISREPRESENTATION**

14
15 273. Plaintiff incorporates by reference each and every preceding paragraph as though fully
16 set forth herein.

17 274. The Defendants falsely and fraudulently have represented and continue to represent to
18 the medical and healthcare community, Plaintiff and her Prescribing and Administering Health Care
19 Providers, and the public in general that Depo-Provera has been appropriately tested and was found to
20 be safe and effective.
21

22 275. At all times material herein, Defendants misrepresented to consumers and physicians,
23 including Plaintiff and Plaintiff's physicians and the public in general, that Depo-Provera is safe for
24 use as a contraceptive and for other indications.
25

26 276. Defendants knew or should have known of the falsity of such a representation to
27 consumers, physicians, and the public in general since Depo-Provera is far from the only contraceptive
28

1 approved by the FDA, and it is not the only contraception option. Nevertheless, Defendants' marketing
2 of Depo-Provera falsely represented Depo-Provera to be a safe and effective contraceptive option with
3 no increased risk of intracranial meningioma and sequelae related thereto.

4 277. The representations were, in fact, false. When the Defendants made these
5 representations, it knew and/or had reason to know that those representations were false, and
6 Defendants willfully, wantonly, and recklessly disregarded the inaccuracies in their representations
7 and the dangers and health risks to users of Depo-Provera.

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9 278. Prior to Plaintiff's use of Depo-Provera, Defendants knew or should have known of
10 adverse event reports indicating the development of intracranial meningioma in individuals who had
11 taken Depo-Provera.

12
13 279. These representations were made by the Defendants with the intent of defrauding and
14 deceiving the medical community, Plaintiff, and the public, and also inducing the medical community,
15 Plaintiff, Plaintiff's Prescribing and Administering Health Care Providers, and/or the public, to
16 recommend, prescribe, dispense, and purchase Depo-Provera for use as a contraceptive and other
17 treatment indications while concealing the drug's known propensity to cause serious and debilitating
18 intracranial meningioma and sequelae related thereto.

19
20 280. Despite the fact that the Defendants knew or should have known of Depo-Provera's
21 propensity to cause serious and potentially debilitating injuries due to the development of intracranial
22 meningioma and sequelae related thereto, the label did not contain any of this information in the
23 "Warnings" section. In fact, the label for Depo-Provera has been updated at least a dozen times over
24 the past 20 years, yet at no point did Defendants provide any of the foregoing information in the
25 "Warnings" section. To date, the Depo-Provera label still does not include any warnings whatsoever
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1 that indicate the dangers of intracranial meningioma and sequela related thereto after using Depo-
2 Provera.

3 281. In representations to Plaintiff and/or to her healthcare providers, including Plaintiff's
4 prescribing physician, the Defendants fraudulently stated that Depo-Provera was safe and omitted
5 warnings related to intracranial meningioma.
6

7 282. In representations to Plaintiff and/or to her Prescribing and Administering Health Care
8 Providers, Defendants fraudulently stated that Depo-Provera was safe and concealed and intentionally
9 omitted material information from the Depo-Provera product labeling in existence at the time Plaintiff
10 was prescribed Depo-Provera in 2005.
11

12 283. Defendants were under a duty to disclose to Plaintiff and her physicians the defective
13 nature of Depo-Provera, including but not limited to, the propensity to cause the development of
14 intracranial meningioma, and consequently, its ability to cause debilitating and permanent injuries.
15

16 284. The Defendants had a duty when disseminating information to the public to disseminate
17 truthful information; and a parallel duty not to deceive the public, Plaintiff, and/or her physicians.
18

19 285. The Defendants knew or had reason to know of the dangerous side effects of Depo-
20 Provera as a result of information from case studies, clinical trials, literature, and adverse event reports
21 available to the Defendants at the time of the development and sale of Depo-Provera, as well as at the
22 time of Plaintiff's prescription.

23 286. Defendants' concealment and omissions of material facts concerning the safety of the
24 Depo-Provera were made purposefully, willfully, wantonly, and/or recklessly to mislead Plaintiff ,
25 Plaintiff's physicians, surgeons and healthcare providers and to induce them to purchase, prescribe,
26 and/or use the drug.
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1 287. At the time these representations were made by Defendants, and at the time Plaintiff
2 and/or her Prescribing and Administering Health Care Providers used Depo-Provera, Plaintiff and/or
3 her Prescribing and Administering Health Care Providers were unaware of the falsehood of these
4 representations.

5 288. In reliance upon these false representations, Plaintiff was induced to, and did use Depo-
6 Provera, thereby causing severe, debilitating, and potentially permanent personal injuries and damages
7 to Plaintiff. The Defendants knew or had reason to know that the Plaintiff had no way to determine
8 the truth behind the Defendants' concealment and omissions, and that these included material
9 omissions of facts surrounding the use of Depo-Provera as described in detail herein.
10

11 289. In comporting with the standard of care for prescribing physicians, Plaintiff's
12 prescribing physicians relied on the labeling for Depo-Provera in existence at the date of prescription
13 that included the aforementioned fraudulent statements and omissions.
14

15 290. These representations made by Defendants were false when made and/or were made
16 with the pretense of actual knowledge when such knowledge did not actually exist, and were made
17 recklessly and without regard to the true facts.
18

19 291. Plaintiff did not discover the true facts about the dangers and serious health and/or
20 safety risks, nor did Plaintiff discover the false representations and omissions of the Defendants, nor
21 could Plaintiff with reasonable diligence have discovered the true facts about the Defendants'
22 misrepresentations at the time when Depo-Provera was prescribed to her.
23

24 292. As a direct and proximate result of reliance upon Defendants' fraudulent
25 misrepresentations, Plaintiff suffered bodily injuries and resulting pain and suffering, disability,
26 mental anguish, loss of capacity for the enjoyment of life, expense of medical and nursing care and
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1 treatment, loss of earnings, loss of consortium, loss of ability to earn money and other economic losses.
2 The losses are either permanent or continuing, and Plaintiff will suffer the losses in the future.

3 293. Defendants have engaged in willful, malicious conduct and/or conduct so careless that
4 it demonstrates a wanton disregard for the safety of others, including Plaintiff, such that the imposition
5 of punitive damages is warranted here.

6 **COUNT VIII**

7 **BREACH OF EXPRESS WARRANTY**

8
9 294. Plaintiff incorporates by reference each and every preceding paragraph as though fully
10 set forth herein.

11 295. At all relevant times herein, Defendants engaged in the business of researching, testing,
12 developing, manufacturing, labeling, marketing, selling, inspecting, handling, storing, distributing,
13 and/or promoting Depo-Provera, and placed it into the stream of commerce in a defective and
14 unreasonably dangerous condition. These actions were under the ultimate control and supervision of
15 Defendants.

16
17 296. Defendants expressly warranted to Plaintiff, Plaintiff's Prescribing and Administering
18 Health Care Providers, and the general public, by and through Defendants and/or their authorized
19 agents or sales representatives, in publications, labeling, the internet, and other communications
20 intended for physicians, patients, Plaintiff, and the general public, that Depo-Provera was safe,
21 effective, fit and proper for its intended use.

22
23 297. Depo-Provera materially failed to conform to those representations made by
24 Defendants, in package inserts and otherwise, concerning the properties and effects of Depo-Provera,
25 which Plaintiff purchased and consumed via intramuscular injection in direct or indirect reliance upon
26 these express representations. Such failures by Defendants constituted a material breach of express
27 warranties made, directly or indirectly, to Plaintiff concerning Depo-Provera as sold to Plaintiff.
28

BREACH OF IMPLIED WARRANTY

305. Plaintiff incorporates by reference each and every preceding paragraph as though fully set forth herein.

306. At all relevant times herein, Defendants engaged in the business of researching, testing, developing, manufacturing, labeling, marketing, selling, inspecting, handling, storing, distributing, and/or promoting Depo-Provera, and placed it into the stream of commerce in a defective and unreasonably dangerous condition. These actions were under the ultimate control and supervision of Defendants.

307. Defendants were the sellers of the Depo-Provera and sold Depo-Provera to be taken for contraception or to treat endometriosis, among other indications. Plaintiff was prescribed and purchased Depo-Provera for these intended purposes.

308. When the Depo-Provera was prescribed by Plaintiff's physicians and taken by Plaintiff, the product was being prescribed and used for the ordinary purpose for which it was intended.

309. Defendants impliedly warranted their Depo-Provera product, which they manufactured and/or distributed and sold, and which Plaintiff purchased and ingested, to be of merchantable quality and fit for the common, ordinary, and intended uses for which the product was sold.

310. Defendants breached their implied warranties of the Depo-Provera product because the Depo-Provera sold to Plaintiff was not fit for its ordinary purpose as a contraceptive or to treat endometriosis safely and effectively, among other uses.

311. The Depo-Provera would not pass without objection in the trade; is not of fair average quality; is not fit for its ordinary purposes for which the product is used; was not adequately contained, packaged and labeled; and fails to conform to the promises or affirmations of fact made on the container or label.

312. Defendants' breach of their implied warranties resulted in the intramuscular

1 administration of the unreasonably dangerous and defective product into Plaintiff, which placed
2 Plaintiff's health and safety at risk and resulted in the damages alleged herein.

3 313. As a direct and proximate result of reliance upon Defendants' breaches of warranty,
4 Plaintiff suffered bodily injuries and resulting pain and suffering, disability, mental anguish, loss of
5 capacity for the enjoyment of life, past and future medical care and treatment, loss of earnings, loss of
6 consortium, loss of ability to earn money and other economic losses, and other damages. The losses
7 are either permanent or continuing, and Plaintiff will suffer the losses in the future.
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PRAYER FOR RELIEF

WHEREFORE, Plaintiff respectfully requests that the Court:

1. Award Plaintiff compensatory and punitive exemplary damages in an amount to be determined at trial, and also including, but not limited to:
 - a. General Damages for severe physical pain, mental suffering, inconvenience, and loss of the enjoyment of life;
 - b. Special Damages, including all expenses, incidental past and future expenses, medical expenses, and loss of earnings and earning capacity;
2. Award interest as permitted by law;
3. Award reasonable attorneys' fees and costs, as provided for by law; and
4. Grant such other and further relief as the Court deems just and proper.

DEMAND FOR JURY TRIAL

Plaintiff demands a trial by jury on all Counts and as to all issues.

Dated: November 4, 2024

Respectfully Submitted,

By: /s/ Melinda Davis Nokes

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